

The stem rust resistance gene *Rpg5* encodes a protein with nucleotide-binding-site, leucine-rich, and protein kinase domains

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We isolated the barley stem rust resistance genes *Rpg5* and *rpg4* by map-based cloning. These genes are colocalized on a 70-kb genomic region that was delimited by recombination. The *Rpg5* gene consists of an unusual structure encoding three typical plant disease resistance protein domains: nucleotide-binding site, leucine-rich repeat, and serine threonine protein kinase. The predicted RPG5 protein has two putative transmembrane sites possibly involved in membrane binding. The gene is expressed at low but detectable levels. Posttranscriptional gene silencing using VIGS resulted in a compatible reaction with a normally incompatible stem rust pathogen. Allele sequencing also validated the candidate *Rpg5* gene. Allele and recombinant sequencing suggested that the probable *rpg4* gene encoded an actin depolymerizing factor-like protein. Involvement of actin depolymerizing factor genes in nonhost resistance has been documented, but discovery of their role in gene-for-gene interaction would be novel and needs to be further substantiated.

actin depolymerizing factor | barley | disease resistance domains | map-based cloning

Stem rust caused by the fungus *Puccinia graminis* (*Pg*) was historically one of the most significant foliar diseases of barley and wheat, with genetic resistance being the primary means of control. Durable resistance in barley has been achieved against many pathotypes for the past 60+ years by the widespread use of the single resistance gene, *Rpg1* (1).

Barley can be attacked by *Pg* f. sp. *tritici* (*Pgt*), the wheat stem rust pathogen and *Pg* f. sp. *secalis* (*Pgs*), the rye stem rust pathogen. In cultivated barley, five genes are known to confer resistance to *Pgt* and three to *Pgs* (2, 3). Only *Rpg1* has been cloned and characterized (4, 5). A virulent *Pgt* pathotype, designated QCC, was isolated from Midwestern barley cvs. containing *Rpg1* in 1989 (6). A resistance gene identified in barley line Q21861 was designated *rpg4*. It acts in recessive manner to pathotype QCC and is temperature-sensitive (3, 7, 8). Besides *rpg4*, Q21861 carries *Rpg5* providing resistance to *Pgs* isolate 92-MN-90 (3, 8). The *Rpg5* gene, previously *RpgQ*, is dominant or semidominant in action and was reported to cosegregate with *rpg4*, although three exceptions were found in 769 F₂ progeny (9). The *rpg4* gene was mapped to the long arm of barley chromosome 7(5H) (10). More detailed mapping and identification of syntenic rice chromosome regions (11, 12) allowed the development of a physical map covering the presumed *rpg4* gene region (13).

Advances in molecular techniques and tools have facilitated the cloning of numerous plant disease resistance genes (*R* genes) in the past two decades. *R* genes are grouped into different

classes according to their protein domain structure (14). The largest group consists of the NBS-LRR family of *R* genes, which is characterized by an N-terminal nucleotide-binding site (NBS) and C-terminal leucine rich repeats (LRRs). Another class, with relatively few members, is the extracellular LRR and transmembrane (TM) domain containing genes conferring resistance to the fungus *Cladosporium fulvum*, the leaf mold pathogen of tomato. The rice genes *Xa21* (15) and *Xa26* (16) conferring resistance to the bacterial blight pathogen *Xanthomonas oryzae* are the only examples of receptor-like kinase genes consisting of an extracellular LRR, a TM, and a cytoplasmic serine/threonine protein kinase (S/TPK). Last, there is a class of *R* genes that consists of S/TPK domains. S/TPK *R* genes include the previously described barley *Rpg1* gene (4); the tomato *Pto* gene, which confers resistance to the bacterial pathogen *Pseudomonas syringae* pv. *tomato* (17); and the *Arabidopsis* *PBS1* gene, which confers resistance to the bacterial pathogen *Pseudomonas syringae* pv. *phaseolicola* (18).

The S/TPK group of *R* genes is unique in that two of the members, *Pto* and *PBS1*, have been shown to require an NBS-LRR gene, *Prf* and *RPS5*, respectively, for resistance (19, 20). This demonstrates that NBS-LRR and protein kinases sometimes work together to provide resistance to plant pathogenic organisms.

Here, we report the cloning and preliminary characterization of two unique barley stem rust resistance genes *Rpg5* and *rpg4*. *Rpg5* encodes an *R* gene protein containing the NBS, LRR, and S/TPK domains in a single transcript. We validated the candidate *Rpg5* gene by multiple allele sequencing and VIGS (21).

Available recombinants point to the *rpg4* gene encoding an actin depolymerizing factor (Adf). Adfs play an important role

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The authors declare no conflict of interest.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database [accession nos. EU883792 (*Rpg5* cDNA from barley line Q21861); EU878778 (genomic sequence from barley line Q21861); EU812563 (genomic sequence from barley line Morex); EU881932-EU881935 (*HvAdf2* genomic sequence from barley lines Golden Promise, Harrington, MD2, and Steptoe, respectively); EU883581-EU883583 (*HvRGA1* genomic sequence from barley lines Harrington, MD2 and Steptoe, respectively); EU883787-EU883790 (*HvRGA2* genomic sequence from barley lines Golden Promise, Harrington, MD2, and Steptoe, respectively); and EU883791 (*HvAdf3* genomic sequence from barley line Steptoe)].

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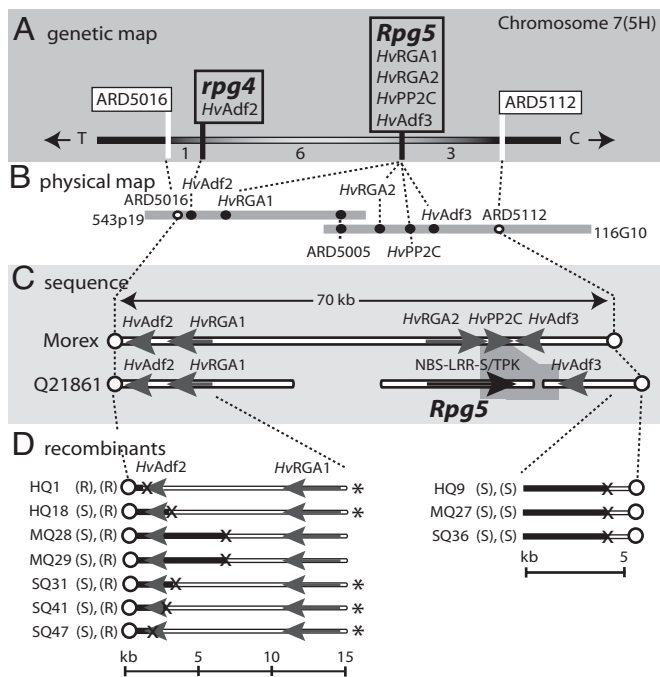


Fig. 1. Genetic, physical, sequence, and recombinant characterization of the *Rpg5/rpg4* locus. (A) High-resolution genetic map of the *Rpg5* region based on 5,232 recombinant gametes. The *Rpg5* flanking markers (ARD5016 and ARD5112) are designated above the white vertical bars. The candidate genes cosegregating with isolate 92-MN-90 (*Rpg5*) resistance and pathotype QCC (*rpg4*) resistance are inside the boxes labeled *Rpg5* and *rpg4*, respectively. Numbers below the bar indicate the number of cross-overs. C and T represent centromere and telomere. (B) BAC physical map spanning the *Rpg5/rpg4* locus. The two BAC clones (543P19 and 116G10) spanning the region are shown as horizontal gray bars. The white circles indicate the flanking genetic markers. Black circles indicate the position of *Rpg5* and *rpg4* candidate genes and the marker (ARD5005) used for chromosome walking. (C) Sequence annotation of the *Rpg5* region. White horizontal bars represent sequenced regions from Morex and Q21861. White circles show the position of flanking markers. The black arrow represents the *Rpg5* NBS-LRR-S/TPK gene. Gray arrows indicate annotated genes. The darker gray background between *HvRGA2* and *HvAdf3* indicates the region of colinearity breakdown. The scale is shown above in kilobases. (D) Recombinant sequence analysis. Horizontal bars represent sequence and/or genotyping from 10 lines with recombination defining the *Rpg5/rpg4* region. Black indicates susceptible genotype, and white indicates the resistant Q21861 genotype. The "X" indicates the approximate point of recombination. The recombinant designations are labeled to the right with resistant (R) or susceptible (S), indicating the response to pathotype QCC (*rpg4*) and isolate 92-MN-90 (*Rpg5*), respectively. Gray arrows indicate candidate genes and the white circles represent flanking markers. Recombinant lines indicated with an asterisk have complete sequence analysis at the region of recombination. The scale is shown below in kilobases.

in cytoskeleton organization and have been reported to be involved with nonhost disease resistance (22). Their role in gene-for-gene interaction would be novel and needs to be further substantiated.

Results

High-Resolution Genetic and Physical Mapping. High-resolution genetic mapping identified the *Rpg5/rpg4* containing region between RFLP markers ARD5112 and ARD5016. This region was delimited by 10 cross-overs, 7 in the *HvAdf2* region and 3 between ARD5112 and *HvAdf3* (Fig. 1 A and D). The recombinants around the *HvAdf2* region separated *rpg4* (resistance to *Pgt* pathotype QCC) from *Rpg5* (resistance to *Pgs* isolate 92-MN-90) (Fig. 1D).

Physical mapping identified BAC clones 543P19 and 116G10

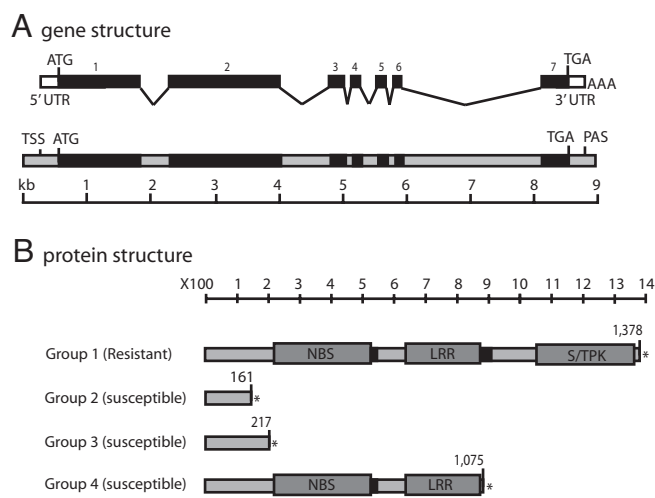


Fig. 2. *Rpg5* predicted gene structure from resistant and susceptible cultivars. (A) cDNA and genomic DNA structures with introns (gray), exons (black), and UTRs (white) shown to scale. *Rpg5* predicted mRNA structure is shown above the genomic sequence with exons numbered above. ATG represents the start methionine codon and TGA represents the stop codon. AAA is the mRNA poly(A) tail, TSS is the transcription start site, and PAS is the polyadenylation signal. The scale below is shown in kilobases. (B) Organization of the predicted protein structure is shown to scale with predicted boundaries indicated by boxes. NBS denotes the nucleotide-binding site, LRR denotes 12 imperfect leucine-rich repeats, S/TPK denotes the serine threonine protein kinase domain, and the black bars represent predicted transmembrane domains. Numbers above indicate the last amino acid preceding the stop codon. Stop codons are represented by an asterisk. Cultivars or lines were placed into four groups. Group 1 (resistant) consists of Q21861 with a predicted intact and functional *Rpg5* gene. Group 2 (susceptible) consists of the cultivars Morex and Steptoe. They have a cytosine-to-adenine conversion that introduces a stop codon resulting in a predicted truncated protein. In addition, they are missing the kinase domain coding region. Group 3 (susceptible) contains Golden Promise and MD2. They have a single cytosine insertion causing a frame-shift mutation that results in a stop codon and a predicted truncated protein. Group 4 (susceptible) consists of Harrington. Harrington is missing the kinase coding region and the second transmembrane domain.

spanning the ARD5112 to ARD5016 region (Fig. 1B). Genetic and physical mapping confirmed that these BAC clones included the flanking markers and the *Rpg5/rpg4* locus.

Identification and Analysis of Candidate *Rpg5* and *rpg4* Genes. Sequence analysis of the cv. Morex BAC clones 543P19 and 116G10 identified a 70-kb region flanked by the RFLP markers ARD5016 and ARD5112. This region was annotated, and five candidate genes were identified, two encoding predicted R-like NBS-LRR proteins (*HvRGA1* and *HvRGA2*), two actin depolymerizing factors (*HvADF2* and *HvADF3*), and a protein phosphatase 2C protein (*HvPP2C*) (Fig. 1C).

PCR primers designed to the cv. Morex candidate genes amplified four of the five genes from the resistant line Q21861. The *HvPP2C* gene could not be amplified with Morex-specific PCR primers, suggesting a diverged gene or an indel event. The Q21861 noncolinearity region was amplified by using PCR chromosome walking (23) and sequenced revealing a S/TPK domain associated with the *Rpg5* candidate gene *HvRGA2* (Fig. 2). The S/TPK domain was absent from susceptible cvs. Morex, Steptoe, and Harrington, suggesting its involvement in *Rpg5*-mediated resistance.

To characterize the noncolinearity region further, a Q21861 λ library was developed and clone RSB762 identified. The RSB762 sequence contained the *Rpg5* gene with 5 kb 5' region and 7.5 kb extending 3' into the noncolinearity region. Additional chromosome walking resulted in 15,708-bp Q21861 sequence. This

completed the noncolinearity region except for a small retrotransposon block, estimated at ≈ 2 kb by restriction mapping, which we could not assemble because of its repetitive nature. The S/TPK domain was the only additional gene found in the Q21861 indel region.

An approximately 18-kb region between *HvRGA1* and *Rpg5* contained a large retrotransposon block in Morex. We were not able to assemble Q21861 contiguous sequence from this region. The partial sequences obtained were all repetitive and/or retrotransposon-like, with no evidence of additional genes. The 57.3-kb total Q21861 sequence identified the same candidate genes as in the susceptible Morex sequence, except that the *HvPP2C* gene was replaced by a protein kinase domain.

Comparison of Morex and Q21861 sequences suggested that *HvRGA2* was the *Rpg5* gene (nucleotide and amino acid numbers given are from the Q21861 *Rpg5* sequence, starting with the first nucleotide in the translation start codon and the first amino acid). The Morex *HvRGA2* consisted of typical NBS-LRR domains and a PP2C gene a short distance downstream (Fig. 1C). The Q21861 *HvRGA2* gene consisted of NBS-LRR-S/TPK domains (Figs. 1C and 2). Furthermore, the susceptible cv. Morex genomic and cDNA sequence revealed a nucleotide substitution at position +483, resulting in a stop codon at amino acid position 161 and a predicted truncated protein (Fig. 2B, Group 2). This correlation of an apparently nonfunctional protein and a susceptible phenotype suggested that *HvRGA2* could be the *Rpg5* gene. Sequence analysis of the *HvRGA2* alleles from the susceptible parents showed Steptoe to be similar to Morex, encoding a predicted truncated protein. MD2 and Golden Promise contained a single-nucleotide insertion (C337) causing a frame-shift mutation at amino acid position 114 resulting in a stop codon at aa position 217 (Fig. 2B, Group 3). The susceptible cv. Harrington *HvRGA2* allele contained an amino acid sequence very similar to Q21861, but it was lacking the PK domain (Fig. 2B).

The remaining candidate genes were eliminated based on allele sequencing. The candidate gene *HvRGA1* codes for an 895-aa (98.5-kDa) predicted NBS-LRR protein with highest homology to an *Oryza sativa* hypothetical NBS-LRR protein (GenBank accession no. EAY80410). Sequence analysis of *HvRGA1* alleles revealed that the resistant parent Q21861 and the susceptible parent MD2 had identical *HvRGA1* predicted amino acid sequences. The susceptible parent Harrington differed from Q21861 by only a single amino acid (C324R). The susceptible cvs. Morex and Steptoe were identical at the amino acid level and differed from Q21861 by five amino acids (S290A, K340N, A445G, N474D, and F586L). The single amino acid difference between Q21861 and Harrington occurred within the NBS domain but was not within one of the highly conserved sub domains. Two of the amino acid differences between Q21861 and Morex/Steptoe occurred within the NBS domain and three occurred within the predicted LRR region. The amino acid sequence identity between Q21861 and MD2 in addition to the very limited polymorphism between Q21861 and Harrington suggested that *HvRGA1* was not *Rpg5*.

The *HvAdf3* gene codes for a protein with the highest amino acid homology to a *Lophopyrum elongatum* actin depolymerizing factor-like gene (GenBank accession no. AAG28460). Allele sequencing from the mapping population parents (Steptoe, MD2, and Q21861) and cv. Morex showed that *HvADF3* had no polymorphism at the amino acid level, suggesting that it is a highly conserved protein and not the *Rpg5* gene.

The *HvPP2C* gene sequence from cv. Morex codes for a protein with the highest homology to an expressed *O. sativa* protein phosphatase 2C family protein (GenBank accession no. ABF99721). Specific *HvPP2C* primers (from Morex) failed to amplify Q21861, MD2 or cv. Golden Promise (susceptible), suggesting that either the *HvPP2C* gene was not present or was

highly diverged. Southern blot analysis confirmed that *HvPP2C* was absent from Q21861 genome (data not shown). Sequence analysis of the λ clone RSB762 confirmed the absence of *HvPP2C* gene from this region of the resistant line Q21861.

Genetic mapping identified seven recombination events between the markers ARD5016 and *HvRGA1*, a physical region of 12 kb containing *HvAdf2* and *HvRGA1* (Fig. 1D). The sites of recombination were identified to within ≈ 200 -bp intervals by sequencing and SNP analysis. Recombinants with the susceptible cvs. *HvAdf2* allele were resistant to the stem rust isolate 92-MN-90, eliminating *HvAdf2* from consideration as the *Rpg5* gene (Fig. 1D). However, the six recombinants differentiating *rpg4* (*Pgt* pathotype QCC resistance) from *Rpg5* (*Pgs* isolate 92-MN-90 resistance) and an additional recombinant occurring distal of *HvAdf2* (HQ1) identified *HvAdf2* as the probable *rpg4* gene.

HvAdf2 codes for a 147 aa (16.2 kDa) actin-depolymerizing factor-like protein with highest homology to the *O. sativa* actin-depolymerizing factor 4 expressed gene (GenBank accession no. ABF99587.1). Sequence analysis of the *HvAdf2* alleles revealed that the resistant parent Q21861 and susceptible parent Steptoe and cv. Morex differed by three amino acids (Q39H, A101T, and S135G). However, the pathotype QCC susceptible parent Harrington had an *HvAdf2* gene identical to the Q21861 allele at the amino acid level.

Structure and Expression of the *HvRGA2* (*Rpg5*) Gene. RT-PCR analysis of *HvRGA2* showed it was expressed at the mRNA level in all parents tested, indicating that transcription did not correlate with resistance. However, PCR primers designed to amplify the junction between the LRR domain and the S/TPK domain produced an RT-PCR product only from Q21861, MD2, and Golden Promise [supporting information (SI) Table S1]. Although MD2 and Golden Promise contain the intact NBS-LRR-S/TPK transcript, both MD2 and Golden Promise alleles were shown to contain a frame-shift mutation within the N-terminal region of the gene resulting in a stop codon at amino acid position 217 and a putative truncated protein (Fig. 2B, Group 3).

To obtain *HvRGA2* (*Rpg5*) transcription start site (TSS) primers were designed from the genomic sequence in 100 bp (Table S1) increments. They were used in RT-PCRs to delimit the TSS to within 100 bp at position -408 to -346 bp.

HvRGA2 (*Rpg5*) encodes an apparently functional NBS-LRR-S/TPK gene containing seven exons transcribed into a predicted 4.4-kb mRNA coding for a 1,378-aa (151.6-kDa) predicted protein (Fig. 2). The transcript size was confirmed by Northern blot analysis, which showed a single hybridizing band at ≈ 4.8 kb (Fig. 3A).

The S/TPK domain contains all nine conserved amino acids (24), suggesting a functional kinase. The kinase domain had the highest homology to an *O. sativa* putative S/TPK (GenBank accession no. EAZ08788) but also showed significant similarity to the known *R* gene *Pto* (36% amino acid identity and 53% amino acid similarity). The *Rpg5* S/TPK domain was very homologous to the unknown function *Rpg1* gene family members ABC1041 and ABC1063 with 60% and 61% amino acid identity, respectively, and 76% amino acid similarity (25).

The NBS-LRR region contained the 4 NBS conserved sub-domains and 12 imperfect LR repeats. This region had the highest homology to an *O. sativa* hypothetical protein (GenBank accession no. EAY98635). The *Rpg5* NBS-LRR is very similar to the rice gene *Pi-ta* (40% amino acid identity and 54% amino acid similarity) conferring resistance to the rice blast fungus (26).

The TMPRED program (www.ch.embnet.org/software/TMPRED_form.html) predicted two transmembrane domains, one on the C-terminal side of the NBS domain and the other on the C-terminal side of the LRR domain (Fig. 2B, Group 1).

barleys (B.S., unpublished data). The observation that the QCC-susceptible cv. Harrington has an expressed *Adf2* gene identical to the resistant Q21861 allele at the amino acid level also suggested that additional factors may be involved. One hypothesis is that the recessive *rpg4* gene functions as a pathotype QCC-specific disease-resistance gene only in the presence of another functional disease resistance gene, perhaps *Rpg5*.

ADFs are critical in remodeling the actin cytoskeleton during normal plant development and under biotic and abiotic stress. They are typically small proteins that function in rapid recycling of actin monomers (28). Actin functions in cytoskeleton organization, which coordinates essentially all aspects of plant growth (29). Actin microfilament polymerization has been shown to be involved in nonhost disease resistance. For example, using cytochalasin E, an inhibitor of actin microfilament polymerization and *Arabidopsis* defense-related mutants *eds1*, *pad4*, and *nah4*, it was shown that nonhost resistance to the wheat powdery mildew fungus *Blumeria graminis* f. sp. *tritici* in *Arabidopsis* largely depends on actin cytoskeleton dynamics and function of the *EDS1* gene (30). Similar observations have been documented in other systems.

The posttranscriptional gene silencing of *Rpg5* by VIGS showed significant reduction of *Rpg5* transcript at both sampling time points 0 d (seedlings pooled) and 14 d (susceptible and resistant plants sampled separately). Approximately 30% of the BSMV-*asRpg5* plants resulted in a susceptible reaction to the rust fungus. At the 14-d time point, the level of silencing observed corresponded to the phenotype observed with the susceptible plants showing significant reduction in the *Rpg5* mRNA, whereas the resistant plants had mRNA level similar to the controls (Fig. 4). The time-course experiments indicated that *Rpg5* mRNA levels were variable over time. However, induction by fungal infection could not be established because of similar observations in mock and uninoculated controls. The variable *Rpg5* mRNA levels were surprising, because previous experiments with *Rpg1* showed low but steady mRNA levels with or without fungal infection (31, 32).

In summary, we have identified two types of plant disease-resistance genes. The *Rpg5* gene combines features of NBS-LRR type disease-resistance gene with the S/TPK domain, suggesting that this gene may function both in pathogen perception and signal transduction. The candidate gene for *rpg4* needs further verification, but, if confirmed, it would show involvement of actin cytoskeleton in race-specific disease resistance.

Materials and Methods

Plant Materials. Progeny from the crosses Steptoe/Q21861, Harrington/Q21861, and MD2/Q21861 were used for genetic mapping. Q21861 is the source of the stem rust-resistance genes *Rpg5* and *rpg4* (3, 9). Steptoe and Harrington are barley cultivars, and MD2 is a genetic stock with multiple dominant mutations. All three are susceptible to stem rust. Plants were grown in the greenhouse with day/night temperatures of 18°C/14°C, respectively. Metal halide lights supplemented a 16-/8-h light/dark photoperiod.

Molecular Markers. RFLP markers were generated as described (13) or by designing PCR primers based on the cv. Morex or line Q21861 sequence (this study). Primers are described in Table S1.

Genetic and Physical Mapping. High-resolution mapping at the *Rpg5/rpg4* locus used 50 recombinants selected from 5,232 gametes between the flanking markers *Aga5* and ABG391. These 50 recombinants were reduced to 10 between the flanking markers ARD5016 and ARD5112. BAC physical maps were generated as described (13).

Sequencing and Sequence Analysis. The cv. Morex BAC clones (543P19 and 116G10) forming a contig across the *Rpg5/rpg4* region were sequenced at the Institute for Genomics Research (TIGR). All other cultivar DNA was sequenced with the BigDye terminator system on an ABI 373 DNA sequencer (Applied Biosystems) at the Laboratory for Biotechnology and Bioanalysis, Washington State University, Pullman. The λ clone RSB762 was subcloned into the NotI site

of pBluescript (Stratagene) and sequenced by using the EZ::TN™<KAN-2> Insertion Kit (Epicentre). All PCR-generated fragments were either directly sequenced or cloned into pGEM-T Easy vector (Promega) and sequenced by using the EZ::TN™<KAN-2> Insertion Kit (Epicentre). For direct sequencing, gel slices were placed in plugged tips (Rainin Instrument) and frozen at -20°C. After complete thawing, the tip was centrifuged at 4,000 × g for 10 min. The eluate was extracted two times with 24:1 chloroform:isoamyl alcohol and precipitated with 0.3 M NaOAc and 2.5 volumes 95% EtOH. Primers and clones are described in Table S1.

Contigs were assembled by using Vector NTI Advance 9.0 contig express (Invitrogen). The cultivar sequence comparisons were done by using Vector NTI alignX and the National Center for Biotechnology Information bl2seq function (www.ncbi.nlm.nih.gov/blast/bl2seq/wblast2.cgi). All sequences were analyzed with the BLASTX and BLASTN algorithms.

Library Construction. High-molecular-mass Q21861 genomic DNA was isolated by using a modified CTAB method (33). Genomic DNA (≈20 μg) was partially digested with 0.1 units of Sau 3AI for 1 hour to yield a majority of 15- to 23-kb fragments. DNA was dephosphorylated with 0.1 units of calf intestine alkaline phosphatase for 30 min and separated in 1.2% low melting agarose. DNA fragments 9–23 kb were excised from the gel and recovered by β -agarase treatment. Genomic fragments were ligated into the Lambda Dash II vector (Stratagene) predigested with BamHI and packaged using the Gigapack III XL packaging extract (Stratagene).

RNA Isolation and Northern Blot. Isolation of RNA, Northern blot analysis, and hybridization were as described (25).

RT-PCR. Approximately 1 μg of total RNA was used for RT-PCRs by using M-MLV Reverse Transcriptase (Promega) under the manufacturer's recommended conditions. RT-PCR fragments were directly sequenced after elution from agarose gels as described for direct sequencing.

3' and 5' RACE. 3' RACE was carried out by using the 3' RACE System (Invitrogen) following the manufacturer's recommended conditions. 5' RACE was carried out by using the FirstChoice RLM-RACE Kit (Ambion) following the manufacturer's recommended conditions and SMART technology (34). The gene specific primers used for RACE are described in Table S1.

VIGS and qRT-PCR. Barley plants for VIGS experiments were grown in the growth chamber in plastic pots with a day/night temperature of 20°C +/- 1°C and 18°C +/- 1°C, respectively. A 20-/4-h light/dark photoperiod was provided by cool fluorescent tubes (525 uE/m²s).

Rpg5 was silenced by using VIGS as described (21). A 314-bp *Rpg5* cDNA fragment (+2,033 to 2,346 bp) was generated by PCR and ligated in antisense (as) orientation into BSMV-VIGS infectious clone (p γ PDS4as) digested with PacI and NotI (21). The BSMV-*asRpg5* construct was cotranscribed with the α and β genomes of the tripartite BSMV virus by using the mMessage mMachine T7 kit (Ambion), and the T7 promoter. RNA was inoculated onto Q21861 barley plants at the two-leaf stage. The seedlings were inoculated with isolate 92-MN-90 urediniospores at 0.025 mg per plant mixed with a talc carrier ≈11–12 days after virus infection. After inoculation, the plants were misted and placed in the dark under high humidity conditions for 22 h, then exposed to light and misted periodically. After 4 h, the misting was stopped, and the leaves were left to dry slowly. When the leaf surfaces were completely dry, plants were moved to the growth chambers at 20°C and 80% relative humidity. The plants were scored for compatibility or incompatibility at 14 days postfungal infection.

The negative control contained a 121-bp antisense fragment of the multi cloning site (MCS) from pBluescript K/S (Stratagene). The MCS sequence did not hybridize to barley genomic DNA at low stringency conditions, indicating no homologous regions in the barley genome. Q21861 and Steptoe were used as the resistant and susceptible virus uninoculated controls. qRT-PCR was performed on Rotor-Gene 2000 thermocycler (Corbet Research) using the QuantiTect SYBR green PCR system (Qiagen). The *Rpg5* primers used are described in Table S1 and GAPDH primers were described (35). Tissue samples for qRT-PCR were collected at 0 or 14 d after fungal infection. Tissue samples for the time course experiment were collected at 0, 1, 3, 5, 8, 11, and 14 days after fungal inoculation.

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